Ulrich SCHWANEBERG, et al.
VESICLE FOR SEPARATING ...
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Susan J. Mack 202-293-7060
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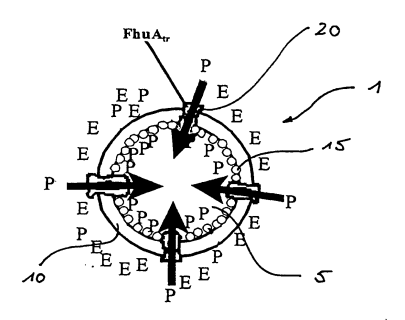


Fig. 1

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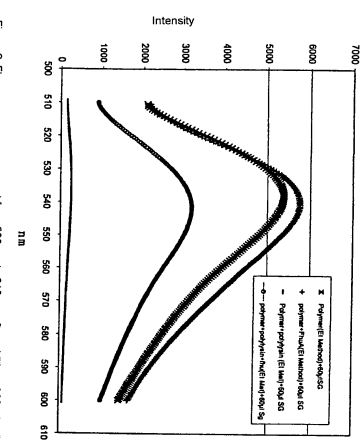
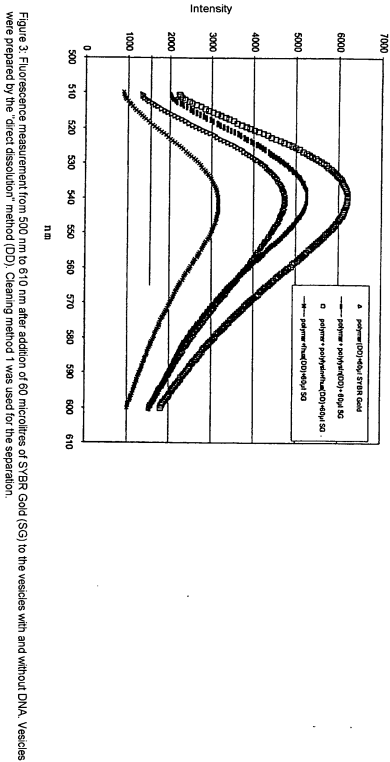
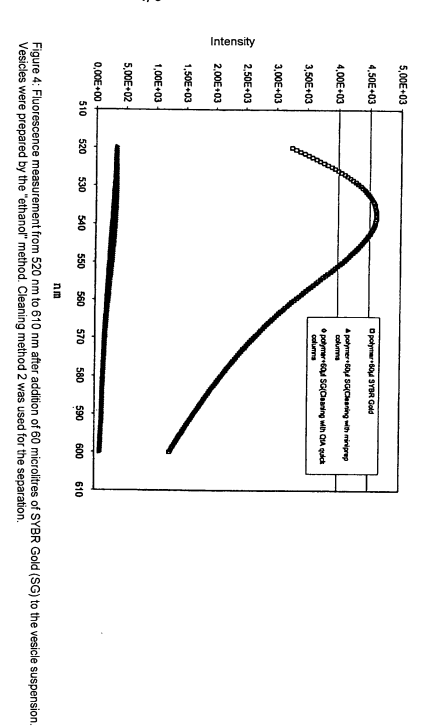


Figure 2: Fluorescence measurement from 500 nm to 610 nm after addition of 60 microlitres of SYBR Gold (SG) to the vesicles with and without DNA. Vesicles were prepared by the "ethanol" method (Et Method; Et Met). Cleaning method 1 was used for the separation.

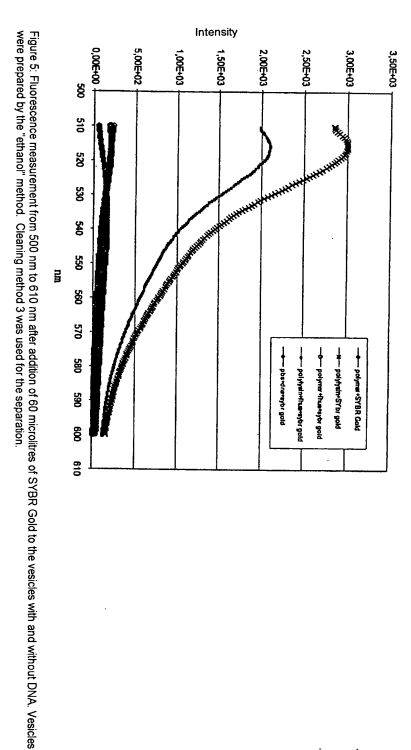
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